A rapid and sensitive one-step test for the qualitative detection of IgG and IgM antibodies to the dengue virus in human serum, plasma, or whole blood. For professional in vitro diagnostic use only.

INTENDED USE
The Accucare Dengue IgG and IgM Combo Rapid Test is a qualitative test for the detection of IgG and IgM antibodies to dengue virus in human serum, plasma or whole blood. The test provides a differential detection of anti-dengue IgG and anti-dengue-IgM antibodies and can be used for the presumptive distinction between a primary and secondary dengue infection. This test is for In-Vitro Diagnostic use only.

SUMMARY
Dengue virus, a virus belonging to the Flaviviridae group of viruses, is one of the most significant mosquito-born diseases in the world in terms of morbidity and mortality. Transmitted principally by the mosquito types Aedes aegypti and Aedes albopictus, the virus is found commonly throughout the tropic and subtropic regions of the world. There are four known serotypes of dengue. Symptoms of dengue fever include high fever, headache, muscle pain and skin rash. The complications often associated with this infection are dengue hemorrhagic fever or dengue shock syndrome. The immune response to this virus includes the production of IgM antibodies by the 5th day of symptoms, which remain in the circulatory system for 30-60 days. IgG antibodies appear by the 14th day of infection and persist for life.

A secondary infection often results in high fever and, in many cases, initiates hemorrhagic events and circulatory failure. A secondary infection also induces an IgM antibody response after 20 days of infection and IgG antibodies rise within 1-2 days after the onset of symptoms. Therefore, patients with secondary infections will have a positive IgG result, usually with a positive IgM result as well. Thus, the use of a reliable and sensitive rapid serological test that can simultaneously detect the presence of anti-dengue IgG and IgM antibodies is of great clinical utility. The Accucare rapid test provides an excellent methodology for specifically detecting anti-dengue IgG and IgM antibodies. The presence of high titers of IgG antibodies does not interfere with the detection of IgM antibodies in the sample. By using a mixture of highly purified dengue proteins, the test is able to detect all 4 Dengue serotypes.

PRINCIPLE
The Accucare Dengue IgG and IgM Combo Rapid Test is a qualitative test for the detection of IgG and IgM antibodies to dengue virus in human serum plasma, or whole blood. The test provides a differential detection of anti-dengue IgG and anti-dengue-IgM antibodies and can be used for the presumptive distinction between a primary and secondary dengue infection. Serum, plasma, or Whole blood samples may be used with this test. First a specimen is dispensed with sample buffer, the latex antigen conjugate will bind to anti-Dengue IgG and IgM antibodies in the specimen sample which in turn will bind with Anti-Human IgG and Anti-Human IgM coated on the membrane as two separate lines in the test region as the reagent move across the membrane. The anti-Human antibodies on the membrane will bind the IgG or IgM antigen complex at the relevant IgG and or IgM test lines causing pale or dark blue lines to form at the IgG or IgM region of the test membrane. The intensity of the lines will vary depending upon the amount of antibody present in the sample. The appearance of blue line in a specific test region (IgG or IgM) should be considered as positive for that particular antibody type (IgG or IgM).

Reagents.
The test strip contains Dengue antigens coated particles and anti-Human IgG and anti-Human IgM coated on the membrane.

PRECAUTIONS
For professional in vitro diagnostic use only. Do not use after the expiration date.

• The test should remain in the sealed pouch until use.
• All specimens should be considered potentially hazardous and handled in the same manner as infectious agents.
• The test should be discarded in a proper biohazard container after testing.
• Optimal assay performance requires strict adherence to the assay procedure described in this Instruction sheet and any deviations from the procedure may lead to aberrant results.

STORAGE & STABILITY
Store as packaged in the sealed pouch at 4 - 30°C and not in direct sunlight. The test is stable through the expiration date printed on the sealed pouch. The test must remain in the sealed pouch until use. DO NOT FREEZE. Do not use beyond the expiration date.

Materials
Materials Provided:
• Test Cassette Device
• Package insert
• Test Buffer
• 1 µL sample loop

Materials Required but not Provided:
• Specimen collection container
• Timer
• Pipette capable of delivering 1 µL sample volume

Specimen Collection & Preparation
• Separate the serum or plasma from blood as soon as possible to avoid hemolysis. Only clear, non-hemolyzed specimens can be used.

Testing should be performed immediately after the specimens have been collected. Do not leave the specimens at room temperature for prolonged periods. Specimens
• may be stored at 2-8°C for up to 3 days. For long-term storage, specimens should be kept below -20°C.
• Bring specimens to room temperature prior to testing. Frozen specimens must be completely thawed and mixed well prior to testing. Specimens should not be frozen and thawed repeatedly.
• If specimens are to be shipped, they should be packed in compliance with federal regulations for the transportation of etiologic agents.

DIRECTIONS FOR USE:
Allow the test device, specimen and/or buffer to equilibrate at room temperature (15-30°C) before testing.

1. Bring the pouch to room temperature before opening it. Remove the test device from the sealed pouch and use it as soon as possible.
2. Place the test device on a clean and level surface. Pipette 1 µL of serum, plasma or whole blood into the sample well (S).
3. Add 4 drops (160 µl) of test buffer to buffer well (A).  
4. Wait for the blue line(s) to appear. The test result should be read at between 15 and 30 minutes. Result may be read up to 60 minutes.  

Note: Do not interpret the result after 60 minutes.

INTERPRETATION OF RESULTS:  

IgM POSITIVE: Two distinct blue lines appear. The control line (C) and IgM (M) line 1 are visible on the test cassette. The test is positive for IgM antibodies. This is indicative of a primary dengue infection.

IgG POSITIVE: Two distinct blue lines appear. The control line (C) and IgG (G) line 2 are visible on the test cassette. The test is positive for IgG antibodies. This is indicative of a secondary dengue infection.

IgM and IgG POSITIVE: Three distinct red lines appear. The control line (C), IgM (M) and IgG (G) lines are visible on the test cassette. The test is positive for IgM and IgG antibodies. This is indicative of a past dengue infection.

NEGATIVE: One distinct blue line appears. The control line (C) is the only line visible on the test cassette. No IgG or IgM antibodies were detected. The result does not exclude dengue infection. A new sample should be drawn from the patient in 3-5 days and then should be retested.

INVALID: Control line fails to appear. The test results are INVALID, if no control line (C) is visible, regardless of the presence or absence of lines in the IgG (G) or IgM (M) region of the cassette. Repeat the test using a new cassette.

NOTE: The intensity of the blue color in the test line regions (G) and (M) will vary depending on the concentration of IgG and IgM present in the specimen. However, neither the quantitative value nor the rate of increase in IgG or IgM can be determined by this qualitative test.

Quality Control  
A procedural control is included in the test. A blue line appearing in the control region (C) is the internal procedural control. It confirms sufficient specimen volume and correct procedural technique. A clear background is also required.

Control standards are not supplied with this kit; however, it is recommended that positive and negative controls be tested as a good laboratory practice to confirm the test procedure and to verify proper test performance.

PERFORMANCE CHARACTERISTICS

<table>
<thead>
<tr>
<th>IgM Results</th>
<th>ELISA Positive</th>
<th>ELISA Negative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Test Positive</td>
<td>41</td>
<td>1</td>
</tr>
<tr>
<td>Rapid Test Negative</td>
<td>1</td>
<td>57</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>IgG Results</th>
<th>ELISA Positive</th>
<th>ELISA Negative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapid Test Positive</td>
<td>40</td>
<td>2</td>
</tr>
<tr>
<td>Rapid Test Negative</td>
<td>2</td>
<td>56</td>
</tr>
</tbody>
</table>

A clinical study using a total of 100 samples was conducted using samples collected in India and Indonesia. The results of the Dengue IgM/IgG Test were compared with a commercially available ELISA test. The sensitivity and specificity of the IgM and IgG test results are given below:

Sensitivity = 97.6%  Specificity = 98.3%
Sensitivity = 95.2%  Specificity = 96.6%

LIMITATIONS AND INTERFERENCES

1. The test procedure, precautions and interpretation of results for the test must be strictly followed.

2. As with all diagnostic tests, the test result must be consistent with clinical findings.

3. Results are to be interpreted within the epidemiological, clinical and therapeutic context.

4. In early primary infections and some secondary infections, detectable levels of IgM antibodies may be low. Some patients may not produce detectable levels of antibody within the first seven to 0 days of infection. If symptoms persist, retest the patient 3 - 5 days after the first testing date.

5. In endemic areas childhood infections with dengue or related flaviviruses are not uncommon and confer life-long levels of IgG antibodies. The sensitivity of the test has been adjusted to minimize detection of very low levels of IgG antibodies, nevertheless, very weak (trace) lines may appear, especially after 30 minutes from the time the test is started. In this case, retest the patient after 5-7 days to look for increased intensity of the IgG line. If there is no change, then dengue is not indicated.

Bibliography


